

(19) World Intellectual Property Organization International Bureau



(43) International Publication Date 17 April 2003 (17.04.2003)

PCT

(10) International Publication Number WO 03/031409 A1

(51) International Patent Classification?: C07D 209/14

(21) International Application Number: PCT/KR02/01843

(22) International Filing Date: 2 October 2002 (02.10.2002)

English

(26) Publication Language:

(25) Filing Language:

English

(30) Priority Data: 2001-0062492 10 October 2001 (10.10.2001) KR

(71) Applicant: CHEIL JEDANG CORPORATION [KR/KR]; 12F, CHEILJEDANG Bldg., 500, 5Ga, Namdaemoon-No, Chung-Ku, Scoul 100-095 (KR).

(72) Inventors: CHO, Il-hwan; 104-102, Hangang Town Apt., Gayang-dong, Gangseo-gu, Seoul 157-200 (KR). LIM, Jee-woong; 960-803, Baikdu-Dongsung Apt., Sanbon-dong, Gunpo Si, Gyeoriggi-Do 435-040 (KR). NOH, Ji-young; 503, Buksan-Green Villa, Jangjeon 2-dong, Geumjeong-gu, Busan 609-392 (KR). KIM, Jong-hoon; 503-503, Gongjak-Lucky Apt., 1587, Gwanyang-dong, Anyang Si Dongan-gu, Gyeonggy-Do 431-060 (KR). PARK, Sang-wook; 201-1505, Woncheon Jugong Apt., 2-Danji, Woncheon-dong, Suwon Si Paldal-gu, Gyeonggi-Do 442-756 (KR). RYU, Hyung-chul; 104, 984-12, Yeongtong-dong, Suwon Si Paldal-gu, Gyeonggi-Do 442-813 (KR). KIM, Je-hak; 110-1403, LG Samik Apt., Homesil-dong, Suwon Si Gwonseon-gu, Gyeonggi-Do 441-708 (KR). CHUN, Hyung-ok; 1132-1204, Samsung-Jangmi Apt., Sanbon-dong, Gunpo Si, Gyeonggi-Do 435-040 (KR). WANG, So-young; 101-212, Joongang-Heights Apt., 305, Cheonho 4-dong,

Gangdong-gu, Seoul 134-867 (KR). LEE, Sung-hak; 766-16 Bangbaebon-dong, Seocho-gu, Seoul 137-829 (KR).

- (74) Agent: CHO, Iu-jae; NEWKOREA INTERNATIONAL PATENT & LAW OFFICE, 3rd Fl., Janghyun Bldg., #637-23, Yeoksam-dong, Gangnam-gu, Seoul 135-909 (KR).
- (81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, UZ, VC, VN, YU, ZA, ZM, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, SK, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

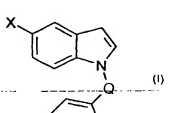
Published:

- with international search report
- before the expiration of the time limit for amending the claims and to be republished in the event of receipt of amendments

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

(54) Title: 1H-INDOLE DERIVATIVES AS A HIGHLY SELECTIVE CYCLOOXYGENASE-2 INHIBITOR

WO 03/031409 A1



(57) Abstract: The present invention relates to a novel 1*H*-indole derivative having a structure of formula (I) and its pharmaceutically acceptable salts as a highly selective cyclooxygenase-2 inhibitor, wherein, X, Y, and Q are defined in this specification respectively.

1H-INDOLE DERIVATIVES AS A HIGHLY SELECTIVE CYCLOOXYGENASE-2 INHIBITOR

TECHNICAL FIELD

5

The present invention relates to 1H-indole derivatives as a highly selective cyclooxygenase-2 inhibitor.

10

15

20

25

BACKGROUND

of non-steroid anti-inflammatory drugs represent actions such as anti-inflammation, ataralgesia, defervescence by inhibiting the enzymatic activity of cyclooxygenase or prostaglandin G/H synthase. In addition, they can suppress the uterine contraction induced by hormones and the proliferation in several kinds of cancers. First, only cyclooxygenase-1 was known to be found in cow as a constitutional enzyme. But recently, cyclooxygenase-2 is elucidated as an induced form. Cyclooxygenase-2 is identified to be discriminated clearly cyclooxygenase-1 and can be provoked easily by mitogen, endotoxin, hormones, growth factors, cytokines and the like.

Prostaglandins have various pathological and physiological functions. Precisely, cyclooxygenase-1 as

a constitutional enzyme participates in the secretion basic endogenous prostaglandin and plays important role in physiological aspects such as stomach homeostasis, renal blood circulation and so on. On the other hand, cyclooxygenase-2 is induced by inflammatory factors, hormones, growth factors, cytokines and the like and thus plays an important role in pathological effects prostaglandins. Therefore, selective inhibitors against cyclooxygenase-2 are expected to have no side effect on account of the functional mechanism compared with the anti-inflammatory drugs such as conventional non-steroid agents represent actions such as anti-inflammation, ataralgesia and defervescence. Furthermore, estimated to suppress the uterine contraction induced by hormones and the cell proliferation in several kinds of cancers. Especially, it probably has lesser side effects such as gastrointestinal toxicity, toxicity and the like. Also, it is assumed to prevent the synthesis of contractive prostanoids and thus inhibit the contraction of smooth muscle induced by the prostanoid. Hence, it can be applied usefully to treat a premature birth, dysmenorrhea, asthma and several diseases associated with eosinophilic leukocytes. Besides, it can be exploited widely to cure osteoporosis, glaucoma and athymia, which has been disclosed in a lot of references, especially the

5

10

15

20

usefulness of selective inhibitors against cyclooxygenase-2 (References: John Vane, "Towards a better aspirin "in Nature, Vol. 367, pp 215-216, 1994; Bruno Battistini, Regina Botting and Y. S. Bakhle, "COX-1 and COX-2; Toward the Development of More Selective NSAIDs "in Drug News and Perspectives, Vol. 7, pp 501-512, 1994; David B. Reitz and Karen Seibert, "Selective Cyclooxygenase Inhibitors "in Annual Reports in Medicinal Chemistry, James A. Bristol, Editor, Vol. 30, pp 179-188, 1995).

The selective inhibitors against cyclooxygenase-2 have been reported to have various structural forms. Among these, the diaryl heterocycle structure, namely a tricyclic system, has been studied most frequently and exploited to construct a lot of candidate substances. In this structure, it is essential that sulfonamide or methanesulfone group exist onto one phenyl group. The initial substance of such a structure is identified to be Dup697 (Bioorganic and Medicinal Chemistry Letters, Vol. 5, No. 18, p 2123, 1995). Then, as a derivative, SC-58635 (Journal of Medicinal Chemistry, Vol. 40, p 1347, 1997) having a pyrrazole structure, MK-966 (WO 95/00501) having a furanone structure and the like are disclosed.

25

5

10

15

purine, teobromine, triethylamine, trimethylamine, tripropylamine, tromethamine and the like.

Besides, the compound of the present invention can be a salt form of nontoxic acids containing the organic acid and the inorganic acid and accepted pharmaceutically, in case that it be basic. Preferably, the acid can be adopted among acetic acid, adipic acid, aspartic acid. 1,5-naphthalenedisulfonic benzenesufonic acid, benzo acid, camposulfonic acid, citric acid, 1,2-ethanedisulfonic acid, ethanesulfonic acid, ethylendiaminetetraacetic acid, fumaric acid, glucoheptonic acid, gluconic acid, glutamic acid, hydriodic acid, hydrobromic acid, hydrochloric acid, icethionic acid, lactic acid, maleic acid, malic acid, manderic acid, methanesulfonic acid, music acid, 2naphthalene disulfonic acid, nitric acid, oxalic acid, parnoic acid, pantothenic acid, phosphoric acid, pivalic acid, propionic acid, salicylic acid, stearic acid, succinic acid, sulfuric acid, tartaric acid, ptoluenesulfonic acid, undecanoic acid, 10-undecenoic acid and the like and more preferably, among succinic acid, hydrobromic acid, hydrochloric acid, maleic acid, methanesulfonic acid, phosphoric acid, sulfuric acid, tartaric acid and the like.

Preferably, the compound of the present invention of formula 1 as a selective inhibitor against cyclooxygenase-2 is that X is NO₂, NH₂, or -

5

10

15

20

NHSO₂CH₃, Y is hydrogen, halogen, C_1 - C_3 -alkyl, or OMe, and Q is C = 0 or CH_2 .

For preferred embodiments of the present invention, the compounds of formula 1 will be described more clearly as follows:

1-benzoyl-5-nitro-1H-indole;

1-benzyl-5-nitro-1H-indole;

1-(4-fluoro-benzyl)-5-nitro-1H-indole;

1-(4-methoxy-benzyl)-5-nitro-1H-indole;

10 1-(4-isopropyl-benzyl)-5-nitro-1H-indole;

1-benzoyl-5-amino-1H-indole;

N-(1-benzyl-1H-indole-5-yl)-methanesulfonamide;

N-[1-(4-fluoro-be:.zyl)-1H-indole-5-yl]-

methanesulfonamide;

N-(1-benzoyl-1H-indole-5-yl)-methanesulfonamide;

1-benzyl-5-nitro-2,3-dihydro-1H-indole;

N-(1-benzyl-2,3-dihydro-1H-indole-5-yl)

methanesulfonamide; and

N-(1-benzoyl-2,3-dihydro-1H-indole-5-yl)-

20 methansulfonamide.

On the other hand, the compounds of formula 1 in the present invention can be prepared by performing the procedures as illustrated below.

25 However, the process for preparing the compounds of the present invention will not be restricted to following descriptions, especially in reaction solvents,

DISCLOSURE OF INVENTION

Based upon the above technical backgrounds, the inventors of the present invention have tried a lot in order to develop novel compounds as a highly selective cyclooxygenase-2 inhibitor. As a result, we have found that 1*H*-indole derivatives of formula 1 satisfied such a purpose and completed the present invention successfully.

Therefore, the object of the present invention is to provide 1*H*-indole derivatives of formula 1 and its pharmaceutically acceptable salts as depicted below.

Hereinafter, the present invention will be described more clearly.

The present invention relates to 1H-indole derivatives of formula 1 and its pharmaceutically acceptable salts.

<Formula 1>

20

5

10

15

Wherein, --- is a double bond or a single bond,

X is NO₂, NH₂, or -NHSO₂R wherein R represents

hydrogen or C_1-C_3 -alkyl,

Y is hydrogen, halogen, C_1-C_3 -alkyl substituted or not substituted by halogen, NO_2 , NH_2 , OH, OMe, CO_2H , or CN.

Q is C=0, C=S, or CH_2 .

The compound of the present invention can exist as a pharmaceutically acceptable salt form, wherein the pharmaceutically acceptable salt means a nontoxic salt containing organic salt and inorganic salt and accepted pharmaceutically. The inorganic salt consists aluminum, ammonium, calcium, copper, iron, lithium, magnesium, manganese, potassium, sodium, zinc and the like and preferabl,, ammonium, calcium, magnesium, organic salt consists of potassium, sodium. The primary-, secondary- or tertiary- amines, naturally substituted amines, cyclic amines, modified salts prepared through a basic ion exchange resin and the like. Preferably, the organic salt can be selected among arginine, betain, caffeine, colin, N, Ndibenzylethylenediamine, diethylamine, 2diethylaminoethanol, 2-dimethylaminoethanol, ethanolamine, ethylenediamine, N-ethylmorpholine, Nethylpiperidine, N-methylglucamine, glucamine, glucosamine, histidine, hydrapamine, N - (2 hydroxyethyl)piperidine, N-(2-hydroxyethyl)pyrrolidine, isopropylamine, lysine, methylglucamine, morpholine, piperazine, piperidine, polyamine resin, procain,

10

15

20

bases, amounts of used reactants and the like.

Moreover, the compound of the present invention also can be prepared by exploiting and combining various synthetic methods described in the present specification or disclosed in other references of those skilled in this arts with a coordinate and arbitrary mode.

Those skilled in the art will appreciate that the conceptions and specific embodiments disclosed in the foregoing description may be readily utilized as a basis for modifying or designing other embodiments.

Concretely, the compound of formula 1 in the present invention can be prepared as illustrated schematically in following reaction formula 1.

5

Wherein, ___, X, Y, and Q are defined above, L is halogen.

As demonstrated in the above reaction formula 1, the compound of the present invention can be prepared through 2 pathways from 5-nitroindole 5nitroindoline as initial material. Namely, benzoyl group or benzyl group are introduced to a nitrogen atom included in the parent nucleus first and then. methanesulfonyl group is inserted to an amine group on 5-location (method 1). On the other methanesulfonyl group is adopted to an amine group on 5-location of parent nucleus first, and then benzoyl group or benzyl group are i..troduced later (method 2).

A detailed description on preparing the compound of the present invention by the above method (1) is as follows.

The reaction of 5-nitroindole or 5-nitroindoline with benzoylhalide or benzylhalide derivatives should be specifically accomplished under the presence of a Concretely, the reaction is performed at the base. range of room temperature 80.C by dimethylformamide. At this moment, the organic base can be selected among triethylamine, trimethylamine, tripropylamine, pyridine, imidazole, and the while the inorganic base can be selected sodiumacetate, sodium hydroxide, sodium hydride, potassium hydroxide, sodium carbonate, potassium

5

10

15

20

carbonate, and the like. More preferably, potassium carbonate can be adopted.

The reaction reducing from nitro to amine is performed under the presence of palladium/ carbon and ammonium formate as a catalyst at the range of room temperature ~ 80°C by using a single or mixed solvent selected among tetrahydrofuran, diethylether, dimethoxyethane, ethylacetate, dichloromethane, methanol, and ethanol.

10 The reaction forming sulfonamide is accomplished as follows: amine and mesyl chloride is reacted under the presence of a base such as triethylamine, trimethylamine, tripropylamine, pyridine, imidazole, and the like by using a solvent selected among 15 tetrahydrofuran, diethylether, dimethoxyethane, ethylacetate, dichloromethane, methanol, ethanol and the like. Preferably, it should be performed at the range of 0 ~ 50°C and more preferably, at a low temperature in between 5 ~ 10°C.

On the other hand, a detailed description on preparing the compound of the present invention by the above method (2) is as follows.

The reduction of 5-nitroindole or 5-nitroindoline as initial material is performed under the same condition with the above method (1) which exploites palladium/ carbon as a catalyst.

Then, the formation of sulfonamide of 5-

25

aminoindole or 5-aminoindoline prepared thereby accomplished as follows: amine and mesyl chloride is reacted under the presence of a base triethylamine, trimethylamine, tripropylamine, pyridine, imidazole, and a solvent selected among tetrahydrofuran, diethylether, dimethoxyethane, ethylacetate, dichloromethane, methanol, and ethanol. Preferably, it should be performed at the range of -30°C ~ room temperature and more preferably, at a low temperature in between -20 ~ -10°C.

The resulting sulfonamide compound will reacted with bezoylhalide or benzylhalide derivatives through the following procedur. The reaction solvent can be a non-reactive solvent such as dichloromethane, diethylether, tetrhydrofuran, and the like. moment, the reaction temperature should be preferably at the range of $-30 \sim 20^{\circ}C$ and more preferably, at a low temperature in between -20 ~ -10°C. A base should be exploited for this reaction, which can be selected among triethylamine, trimethylamine, tripropylamine, pyridine, imidazole and the like as an organic base and among sodium acetate, sodium hydroxide, sodium hydride, potassium hydroxide, sodium carbonate, potassium carbonate and the like as an inorganic base, more preferably sodium hydride.

After completing the reaction, the resulting products can be processed through a common treatment

5

10

15

20

such as chromatography, re-crytallization and the like so as to be separated and purified.

The compound of the present invention depicted in formula 1 has an activity for the selective inhibition against cyclooxygenase-2 and thus can be utilized as an enzymatic inhibitor. The compound of formula 1 having a selective inhibitor against cyclooxygenase-2 can be a substitute conventional non-steroid antifor inflammatory drugs and especially the compound is useful in patients suffering from peptic ulcer, partial enteritis, ulcerative colitis, gastritis, diverticulitis, gastrointestinal haemorrhagia, hypoprothrombinemia and the like as substitute drugs improved in side effects of conventional non-steroid anti-inflammatory drugs. Besides, it is expected to treat inflammatory diseases such as osteoarthritis, rheumatoid arthritis and the like effectively.

The compound of the present invention can be administered in a single dose or in separated doses, depending upon clinical purposes. The specific dosage for patients will vary, depending upon factors such as a sort of drug compound, body weight, sex, physical condition, diet, administration period, administration method, discharge ratio, drug composition and severity of diseases and the like.

The compound of the present invention can be administered as an oral, a local, a parenteral

5

10

15

20

(subcutaneous, venous and muscular silinge or injection), an inhalational or a rectal drug. In case that these are prepared to a pharmaceutical drug, one or more commonly used vehicles, methods for the preparation and the like can be adopted properly from prior arts widely reported to those skilled.

In order to attain the desired purpose of clinical administration, the active compound of formula 1 in the present invention can be administered coincidently by combining more than one component of other commercial drugs.

However, the pharmaceutical drugs containing the compound of the present invention is not limited to forms described above, if it has a purpose for inhibiting cyclooxygenase-2 selectively. All kinds of drugs useful for the enzymatic inhibition can be within the scope of the present invention.

MODES FOR CARRYING OUT THE INVENTION

20

25

15

5

10

Practical and presently preferred embodiments of the present invention are illustrative as shown in the following Examples.

However, it will be appreciated that those skilled in the art, on consideration of this disclosure, may make modifications and improvements within the spirit and scope of the present invention.

<Reference Example 1> Preparation of (1H-indole-5-yl)amine

5 5-nitroindole (1.0g, 6.17 mmol) was dissolved in methanol (10ml) and anhydrous tetrahydrofuran (10ml) at room temperature and then, palladium/ carbon (10%) of a catalystic amount and ammonium formate (2.0g, 31.7 mmol) were added to be stirred slowly at room 10 temperature for 30 minutes. After completing the reaction, the reacting solution was filtered through celite, washed with methanol, concentrated reduced pressure and then, dropped a silica gel short column. Afterward, the residue was concentrated again 15 .. under reduced pressure and triturated with isooctane. As a result, the present compound (0.45g, productive yield 55%) was obtained as a solid phase.

 1 H-NMR (400 MHz, CDCl₃) δ 2.55 (br s, 2H), 6.35 (s, 1H), 6.65 (d, J = 8Hz, 1H), 6.95 (s, 1H), 7.10-7.15 (m, 1H), 7.20 (d, J = 8Hz, 1H), 7.95 (br s, 1H) melting point : 126°C

<Reference Example 2> Preparation of N-(1H-indole-5yl)-methansulfonamide

25

20

(1H-indole-5=yl)-amine (50mg, 0.38mmol) was dissolved in dichloromethane (1.0ml) at -20°C and

trimethyamine (0.063ml, 0.45mmol) and mesyl chlroride (0.032ml, 0.45mmol) were added to be slowly stirred at room temperature for 30 minutes. After completing the reaction, water (5ml) and dichloromethane (5ml) were added additionally and dichloromethane layer was separated. Afterward, the resulting solution was washed with brine, dried over anhydrous magnesium sulfate, concentrated under reduced pressure, and then purified by performing a flash column chromatography (an eluent: ethyl acetate/n-hexane = 1/2, v/v). As a result, the present compound (30mg, productive yield 38%) was obtained.

¹H-NMR (400 MHz, CDCl₃) δ 2.95 s, 3H), 6.25 (br s, 1H), 6.55 (s, 1H), 7.10 (d, J = 8Hz, 1H), 7.25-7.30 (m, 1H), 7.35 (d, J = 8Hz, 1H), 7.5 (s, 1H)

<Reference Example 3> Preparation of N-(2,3-dihydro1H-indole-5-yl)-methansulfonamide

5-nitroindoline (100mg, 0.61mmol) was dissolved in methanol (2ml) and tetrahydrofuran (2ml). Ammonium formate (192mg, 3.05mmol, 5 equivalent) and palladium/carbon (10%) in a catalytic amount were added at room temperature and refluxed at 40°C for ten minutes.

25 After completing the reaction, the reacting solution was filtrated through celite and concentrated under reduced pressure. Afterward, water (5ml) was added to

5

10

the residue, extracted consecutively 4 times with ethyl acetate (10ml), dried over anhydrous magnesium sulfate, concentrated under reduced pressure and then dried completely under a high-degree vacuum. The obtained compound, namely (2,3-dihydro-1H-indole-5-yl)-amine, dissolved in dichloromethane (5ml). trimethylamine (0.063ml, 0.45mmol) was added to be cooled to -20°C and mesyl chloride (0.035ml, 0.45mmol) was added to be stirred for 30 minutes at the same Water (5ml) temperature. was added to separate dichloromethane solution, dried over anhydrous magnesium sulfate, purified through a flash column chromatography (an eluent: ethyl acetate/n-hexane = 2/1, v/v) and then triturated with isooctane. As a result, the present compound (60mg, productive yield 47%) was obtained as a white solid.

 1 H-NMR (400 MHz, DMSO-d₆) δ 2.00 (s, 3H), 3.15 (t, J = 8Hz, 2H), 3.95 (t, J = 8Hz, 2H), 7.25 (d, J = 8Hz, 2H), 7.45-7.60 (m, 4H), 7.80 (s, 1H), 7.95 (d, J = 8Hz, 1H)

<Example 1> Preparation of 1-benzoyl-5-nitro-1H-indole

5-nitroindole (50mg, 0.31mmol) and potassium carbonate (128mg, 0.93mmol) were suspended in dimethylformamide (1.0ml). Then, benzoyl chloride

5

10

(0.04ml,~0.345mmol) was added and stirred at room temperature for 2 hours. After completing the reaction, water and ethylacetate (respectively 5ml) were added to extract, washed with brine, dried over anhydrous magnesium sulfate, concentrated under reduced pressure, and then purified through a flash column chromatography (an eluent: ethyl acetate/n-hexane = 1/1, v/v). As a result, the present compound (35mg, productive yield 43%) was obtained.

10

5

¹H-NMR (400 MHz, CDCl₃) δ 6.80 (d, J = 3Hz, 1H), 7.50 (d, J = 3Hz, 1H), 7.55-7.70 (m, 3H), 7.80 (d, J = 8Hz, 2H), 8.25-8.30 (m, 1H), 8.50 (d, J = 9Hz, 1H), 8.55 (s, 1H)

15

<Example 2> Preparation of 1-benzyl-5-nitro-1H-indole

The reaction was performed through a same method with Example 1, except exploiting benzyl bromide (0.04ml, 0.366mmol) instead of benzoylchloride. As a result, the present compound (40mg, productive yield 51%) was obtained.

 1 H-NMR (400 MHz, CDCl₃) δ 5.35 (s, 2H), 6.75 (t, 25 J = 2Hz, 1H), 7.10-7.15 (m, 2H), 7.25-7.40 (m, 5H), 8.10 (d, J = 9Hz, 1H), 8.65 (d, J = 2Hz, 1H)

melting point : 103 ~ 104°C

<Example 3> Preparation of 1-(4-fluoro-benzyl)-5-nitro1H-indole

The reaction was performed through a same method with Example 1, except exploiting 4-fluorobenzyl bromide (0.04ml, 0.342mmol) instead of benzoylchloride. As a result, the present compound (55mg, productive yield 66%) was obtained.

10

 $^{1}\text{H-NMR}$ (400 MHz, CDCl₃) δ 5.35 (s, 2H), 6.75 (d, J = 3Hz, 1H), 7.00-7.15 (m, 4H), 7.25-7.30 (m, 2H), 8.10 (d, J = 9Hz, 1H), 8.60 (d, J = 2Hz, 1H) melting point : 114 ~ 115°C

15

<Example 4> Preparation of 1-(4-methoxy-benzyl)-5nitro-1H-indole

The reaction was performed through a same 20 method with Example 1, except exploiting 4-methoxybenzyl bromide (0.046ml, 0.339mmol) instead of benzoylchloride. As a result, the present compound (60mg, productive yield 69%) was obtained.

¹H-NMR (400 MHz, CDCl₃) δ 3.80 (s, 3H), 5.30 (s, 2H), 6.70 (d, J = 3Hz, 1H), 6.85 (d, J = 8Hz, 2H), 7.05

(d, J = 8Hz, 2H), 7.25 (d, J = 3Hz, 1H), 7.30 (d, J = 9Hz, 1H), 8.10 (d, J = 9Hz, 1H), 8.60 (s, 1H)

melting point : $110 \sim 111$ °C

5 <Example 5> Preparation of 1-(4-isopropyl-benzyl)-5nitro-1H-indole

The reaction was performed through a same method with Example 1, except exploiting 4-isopropylbenzyl bromide (0.056ml, 0.339mmol) instead of benzoylchloride. As a result, the present compound (65mg, productive yield 72%) was obtained.

 1 H-NMR (400 MHz, CDCl₃) δ 1.20 (s, 3H), 1.25 (s, 3H), 2.90-2.95 (m, 1H), 5.30 (s, 2H), 6.75 (d, J = 3Hz, 1H), 7.05 (d, J = 8Hz, 2H), 7.20 (d, J = 8Hz, 2H), 7.30 (d, J = 3Hz, 1H), 7.35 (d, J = 9Hz, 1H), 8.60 (s, 1H)

melting point : 120 ~ 121°C

20

25

10

<Example 6> Preparation of 1-benzoy1-5-amino-1H-indole

l-benzoyl-5-nitro-1H-indole (50mg) was dissolved in a mixed solvent with methanol (2ml) and tetrahydrofuran (2ml), and ammonium formate of an excess amount and palladium/ carbon (10%) of a catalytic amount were added. The reacted solution was

stirred at around 30°C for 30 minutes to complete the reduction, filtered through celite and then, concentrated under reduced pressure. Afterward, the residue was dissolved again in ethylacetate (10ml), washed with water and brine, dried over anhydrous magnesium sulfate, concentrated under reduced pressure and then, triturated with isooctane and isopropyleter. As a result, the present compound (25mg, productive yield 56%) was obtained as a solid.

10

5

 $^{1}\text{H-NMR}$ (400 MHz, CDCl₃) δ 3.70 (br s, 2H), 6.45 (d, J = 4Hz, 1H), 6.80-6.85 (m, 1H), 6.85 (s, 1H), 7.20 (d, J = 4Hz, 1H), 7.40-7.60 (m, 3H), 7.70 (d, J = 9Hz, 2H), 8.20 (d, J = 9Hz, 1H)

15

1-benzyl-5-nitro-1*H*-indole (50mg, 0.19mmol)

20 was dissolved in tetrahydrofuran (1ml) and methanol (1ml), and ammonium formate of an excess amount and palladium/ carbon (10%) of a catalytic amount were added. The solution was stirred at around 30°C for 30 minutes to complete the reduction, filtrated through celite, concentrated under reduced pressure, dissolved again in dichloromethane (10ml), and then washed with water and brine. After drying the resulting solution

over anhydrous magnesium sulfate, dichloromethane solution containing amine compound (1-benzyl-5-amino-1H-indole) was obtained. Mesyl chloride (0.015ml, 0.19mmol) and triethylamine (0.028ml, 0.20mmol) were added to the above obtained solution and stirred at room temperature for 2 hours to complete the reaction. After adding 2N-hydrochloric acid solution (10ml) to separate layers, dichloromethane solution was induced, washed by using water and brine, dried over anhydrous magnesium sulfate, concentrated under reduced pressure, and then triturated with isooctane and isopropyleter. As a result, the present compound (34mg, productive yield 34%) was obtained as a solid.

15 $^{1}\text{H-NMR}$ (400 MHz, CDCl₃) δ 2.95 (s, 3H), 5.30 (s, 2H), 6.25 (s, 1H), 6.55 (s, 1H), 7.00-7.35 (m, 8H), 7.55 (s, 1H)

Mass (FAB) 300.0 (M +), 601.1 (2M + 1) melting point : 153 ~ 154°C

20

5

10

<Example 8> Preparation of N-[1-(4-fluoro-benzyl)-1Hindole-5-yl]-methansulfonamide

1-(4-fluoro-benzyl)-5-nitro-1H-indole (50mg,

0.16mmol) was dissolved in tetrahydrofuran (1ml) and

methanol (1ml), and ammonium formate of an excess

amount and palladium/ carbon (10%) of a catalytic

amount were added. The solution was stirred at around 30°C for 1 hour to complete the reduction, filtered through celite, concentrated under reduced pressure, dissolved again in dichloromethane (10ml), and then washed with water and brine. After drying the resulting solution over anhydrous magnesium sulfate, dichloromethane solution containing amine compound (1-(4-fluoro-benzyl)-5-amino-1H-indole) obtained. was Mesyl chloride (0.012ml, 0.16mmol) and triethylamine 10 (0.022ml, 0.16mmol) were added to the above obtained solution and stirred at room temperature for 2 hours to complete the reaction. After adding 2N-hydrochloric acid solution (10ml) to separate dichloromethane solution was induced, washed with water 15 and brine, dried over anhydrous magnesium sulfate, concentrated under reduced pressure, and then triturated with isooctane and isopropyleter. result, the present compound (30mg, productive yield 51%) was obtained as a solid.

20

 1 H-NMR (400 MHz, CDCl₃) δ 2.95 (s, 3H), 5.25 (s, 2H), 6.35 (s, 1H), 6.50-6.55 (m, 1H), 6.95-7.25 (m, 7H), 7.55 (s, 1H)

melting point : 96 ~ 97°C

25

<Example 9> Preparation of N-(1-benzoy1-1H-indole-5y1)-methanesulfonamide

1-benzoyl-5-amino-1H-indole (25mg, 0.106mmol) dissolved in dichloromethane (1.0ml) at temperature and mesyl chloride (0.01ml. 0.116mmol) and 5 triethylamine (0.016ml, 0.115mmol) were added. solution was stirred at room temperature for 30 minutes to complete the reducing reaction. After pouring water (2ml), dichloromethane layer was separated, washed with brine, dried over anhydrous magnesium sulfate and then, 10 purified through flash column chromatography (an eluent: ethyl acetate/n-hexane = 1/2, v/v). As a result, the present compound (20mg, productive yield 60%) was obtained.

15 1 H-NMR (400 MHz, CDCl₃) δ 3.00 (s, 3H), 6.40 (s, 1H), 6.60 (d, J = 4Hz, 1H), 7.20 (d, J = 8Hz, 1H), 7.35 (d, J = 4Hz, 1H), 7.50-7.65 (m, 4H), 7.75 (d, J = 8Hz, 2H), 8.40 (d, J = 8Hz, 1H) Mass (FAB) 314 (M +), 629 (2M + 1) melting point : 123 ~ 125°C

<Example 10> Preparation of 1-benzyl-5-nitro-2,3dihydro-1H-indole

Under the presence of nitrogen, 5-nitroindolin (50mg, 0.30mmol) was dissolved in dimethylformamide (2ml) at room temperature and benzyl bromide (0.04ml.

potassium carbonate (0.126ml,3.0 0.34mmol) and equivalent) were added and stirred at room temperature for 48 hours. After completing the reaction, water and ethyl acetate (respectively 5ml) were added to separate layers, washed with brine, dried over anhydrous magnesium sulfate, and concentrated. Finally, the through flash residue was purified chromatography (an eluent: ethyl acetate/n-hexane = 1/4, As a result, the present compound (45mg, productive yield 58%) was obtained.

 $^{1}\text{H-NMR}$ (400 MHz, CDCl₃) δ 3.10 (t, J = 9Hz, 2H), 3.65 (t, J = 9Hz, 2H), 4.45 (s, 2H), 6.35 (d, J = 9Hz, 1H), 7.25-7.45 (m, 5H), 7.90 (s, 1H), 8.05-8.10 (m, 1H) melting point : 73 ~ 74°C

<Example 11> Preparation of N-(1-benzyl-2,3-dihydro-1Hindole-5-yl)-methanesulfonamide

1-benzyl-5-nitro-2,3-dihydro-1H-indole (100mg, 0.39mmol) was dissolved in tetrahydrofuran (1ml) and methanol (1ml), and ammonium formate (124mg, 1.96mmol, 5 equivalent) and palladium carbon (10%) of a catalytic amount were added. Then, the solution was stirred at 40°C for 10 minutes to complete the reduction. After completing the reaction, the solution was filtered through celite, concentrated under reduced pressure,

5

10

dissolved again in ethyl acetate (10ml), washed with water and salt solution, dried over anhydrous magnesium sulfate, concentrated again under reduced pressure and then, dried completely under a high-degree vacuum. 5 Afterward, the obtained residue was dissolved with dichloromethane (2.0ml), cooled to 0°C, blended with triethylamine (0.055ml, 0.39mmol) and mesyl chloride (0.031ml, 0.40mmol) and then, stirred at the same temperature for 30 minutes to complete the reaction. 10 After pouring water (2.0ml) again at room temperature, dichloromethane layer was separated, washed with brine, dried over anhydrous magnesium sulfate, concentrated. Finally, the residue was purified through flash column chromatography (an eluent: ethyl 15 acetate/n-hexane = 1/2, v/v). As a result, the present compound (90mg, productive yield 76%) was obtained.

 1 H-NMR (400 MHz, CDCl₃) δ 2.95 (t, J = 8Hz, 2H), 3.35 (t, J = 8Hz, 2H), 4.25 (s, 2H), 6.00 (s, 1H), 6.40 (d, J = 8Hz, 1H), 6.90 (d, J = 8Hz, 1H), 7.05 (s, 1H), 7.20-7.35 (m, 5H)

Mass (FAB) 302(M +), 605(2M + 1) melting point : $133 \sim 134$ °C

25 <Example 12> N-(1-benzoyl-2,3-dihydro-1H-indole-5-yl)methanesulfonamide

HR:

N-(2, 3-dihydro-1H-indole-5-y1) -

methanesulfonamide (20mg, 0.095mmol) was dissolved in anhydrous dichloromethane (3ml), sodium hydride (0.010g, 50% oil) in was added and then, 5 benzolychloride (0.011ml, 0.095mmol) was added at a temperature under -20°C. The resulting solution was stirred at the same temperature for 1 hour, stirred again at room temperature for 24 hours to complete the reaction. After pouring water **10** . dichloromethane layer was separated, washed with brine, dried over anhydrous magnesium sulfate and then, concentrated under reduced pressure. Afterward, residue was purified through flash column chromatography (an eluent: ethyl acetate/n-hexane = 1/1, v/v) and triturated with isooctane. As a result, the present compound (15mg, productive yield 50%) was obtained as a solid.

 $^{1}\text{H-NMR}$ (400 MHz, CDCl₃) δ 2.90 (s, 3H), 3.20 (t, 20 J = 8Hz, 2H), 4.00 (t, J = 8Hz, 2H), 7.20 (d, J = 8Hz, 1H), 7.40 (d, J = 8Hz, 1H), 7.45-7.60 (m, 3H), 7.80 (m, 2H), 7.90 (d, J = 8Hz, 2H) Mass (FAB) 317.1(M + 1)

25 <Experimental Example> The activity of selective inhibition against cyclooxygenase-2

(1) Experimental procedure

In order to investigate the activity of the present compound for the selective inhibition against cyclooxygenase-2 enzyme pharmacologically, the enzymatic activities inhibiting cyclooxygenase-1 and cyclooxygenase-2 were measured quantitatively.

First of all, the cyclooxygenase-1 was examined through the following procedure.

Peritoneal fluid in which macrophages suspended was extracted from a mouse peritoneal cavity and centrifuged at 4°C, 1,000 rpm for 2 minutes. Then, the supernatant was removed, suspended with 20ml of incomplete RPMI medium [PC/SM (penicilin/streptomycin)] and again centrifuged under the same condition. In addition, the reactant was washed twice and then the cell pellet was suspended with 10 ml of incomplete RPMI 1640 medium so as to prepare a cell suspension. Then, the cell number was calculated with the hemocytometer and adjusted to reach 1 X 106 cells/ml of cell concentration in the final cell suspension. 100µl of the resulting suspension was transferred into each well of 96-well plate and left at 37°C in 5% CO2 with the incubator for about 2 hours in order to attach macrophages. The attached macrophage was washed twice by using PBS buffer, treated to experimental samples in a proper concentration and then blended with 3% FBS-

5

10

15

20

PCT/KR02/01843

RPMI 1640 medium so as to adjust the total volume reaching 200 μ l. The resulting cell was cultivated in the incubator at 37°C in 5% CO₂ for about 12 ~ 16 hours. Then, arachidonic acid was added, adjusting to 10 μ M of a final concentration and incubated at 37°C for more 10 minutes and the supernatant of the reacted solution (~ 180 μ l) was recovered to finish the reaction. In order to quantitate the amount of PGE2 in the samples, the ELISA method recommended from Cayman Chemical company was exploited and the obtained results was used to estimate the inhibition ratio (%) of each compound against cyclooxygenase-1.

Second, the cyclooxygenase-2 was examined through the following procedure.

Peritoneal fluid suspended with macrophages was extracted from a mouse peritoneal cavity centrifuged at 4°C, 1,000 rpm for 2 minutes. Then, the supernatant was removed, suspended using incomplete RPMI medium [PC/SM (penicilin/streptomycin)] and again centrifuged under the same condition. In addition, the reactant was washed twice and then the cell pellet was suspended with 10 ml of incomplete RPMI 1640 medium so as to prepare a cell suspension. Then, the cell number was calculated with the hemocytometer and adjusted to reach 1 \times 10⁶ cells/ml of cell concentration in the final cell suspension. The resulting suspension was treated with aspirin, adjusting 500 µM of final

5

10

15

20

concentration and transderred into each well of 96-well plate in 100 µl respectively. Again, it was left at 37°C in 5% CO2 in the incubator for about 2 hours in order to attach macrophages. The attached macrophage was washed twice by using PBS buffer, treated to experimental samples in a proper concentration and then blended with 3% FBS-RPMI 1640 medium containing 10 μg/ml of LPS in each well. The resulting cell was cultivated in the incubator at 37° C in 5% CO₂ for about 12 ~ 16 hours. Then, arachidonic acid was added, adjusting to 10 µM of a final concentration incubated at 37°C for more 10 minutes supernatant of the reacted solution (~ 180 μ l) was recovered to finish the reaction. In order to quantitate the amount of PGE2 in the samples, the ELISA method recommended from Cayman Chemical company was exploited and the obtained results was used to estimate the inhibition ratio (%) of each compound against cyclooxygenase-2.

20

5

10

15

(2) Experimental results

The experimental results were demonstrated in Table 1 as follows.

PCT/KR02/01843

<Table 1>
Inhibitory effects of cyclooxygenase (COX) (unit: %
inhibition)

Examples	COX-1			COX-2		
Concentration	30 рм	10 μΜ	3 им	300 nM	100 nM	Mn 06
SC-58635 (standard substance)	81.3	66.5	64.3	73.0	59.9	51.2
1	45.8	40.7	33.2	~ 0	~ 0	~ 0
. 2	80.4	68.7	56.7	.22.0	20.7	15.7
3	74.6	64.4	60.4	70.2	58.8	50.1
4	80.1	71.1	60.3	81.5	69.9	55.4
5	54.3	47.1	39.9	61.4	55.4	51.2
6	64.8	57.3	52.3	54.9	46.6	33.4
7	56.4	44.1	30.0	76.8	70.6	59.8
8	53.9	32.3	7.6	29.6	28.6	22.1
9	42.1	31.1	22.8	30.1	25.5	20.4

5

10

In vitro experiments were observed to measure the inhibitional ratios against cyclooxygenase-1 (COX-1) and cyclooxygenase-2 (COX-2). Consequently, in case of the compound of Example 7, N-(1-benzyl-1H-indole-5-yl)-methanesulfonamide, the inhibition effect against cyclooxygenase-2 was identified to be more excellent than a comparative substance and coincidently, the inhibition effect against cyclooxygenase-1 be in much lower level than a comparative substance. That is to

¹⁵ say, the selectivity of cyclooxygenase-2 is confirmed

to be better than any other substances, which proves the structural efficacy of 1H-indole derivatives in the present invention.

5 INDUSTRIAL APPLICABILITY

As demonstrated and confirmed above, the novel compound of 1H-indole derivative is a substitute drug improved in side effects of conventional non-steroids anti-inflammatory drug and is useful for patients suffering from peptic ulcer, gastritis, partial enteritis, ulcerative colitis, diverticulitis, gastrointestinal haemorrhagia, hypoprothrombinemia and the like. Besides, it is expected to treat inflammatory diseases suca as osteoarthritis, rheumatoid arthritis and the like effectively.

Those skilled in the art will appreciate that the conceptions and specific embodiments disclosed in the foregoing description may be readily utilized as a basis for modifying or designing other embodiments for carrying out the same purposes of the present invention.

Those skilled in the art will also appreciate that such equivalent embodiments do not depart from the spirit and scope of the invention as set forth in the appended claims.

10

15

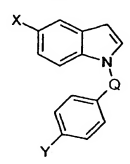
20

5

What is claimed is:

1. A compound of formula 1 and its pharmaceutically acceptable salts:

<Formula 1>



Wherein, -- is a double bond or a single bond, X is NO₂, NH₂, or -NHSO₂R wherein R represents hydrogen or C₁-C₃-alkyl,

Y is hydrogen, halogen, C_1-C_3 -alkyl substituted or not substituted by halogen, NO_2 , NH_2 , OH, OMe, CO_2H , or CN,

Q is C=0, C=S, or CH_2 .

- 15 2. The compound of formula 1 according to claim 1, wherein X is NO_2 , NH_2 , or $-NHSO_2CH_3$, Y is hydrogen, halogen, C_1-C_3 -alkyl, or OMe, and Q is C=O or CH_2 .
- 3. The compound according to claim 1, wherein said compound of formula 1 is selected from a group consisting of:

```
1-benzoyl-5-nitro-1H-indole;
             1-benzyl-5-nitro-1H-indole;
            1-(4-fluoro-benzyl)-5-nitro-1H-indole;
            1-(4-methoxy-benzyl)-5-nitro-1H-indole;
 5
            1-(4-isopropyl-benzyl)-5-nitro-1H-indole;
            1-benzoyl-5-amino-1H-indole;
            N-(1-benzyl-1H-indole-5-yl)-methanesulfonamide;
            N-[1-(4-fluoro-benzyl)-1H-indole-5-yl]-
     methanesulfonamide;
10
            N-(1-benzoyl-1H-indole-5-yl)-methanesulfonamide;
            1-benzyl-5-nitro-2,3-dihydro-1H-indole;
            N-(1-benzyl-2,3-dihydro-1H-indole-5-yl)
     methanesulfonamide; and
            N-(1-benzoyl-2,3-dihydro-1H-indole-5-yl)-
15
     methansulfonamide.
```

INTERNATIONAL SEARCH REPORT

ational application No. PCT/KR02/01843

A. CLA	A. CLASSIFICATION OF SUBJECT MATTER							
IPC?	IPC7 C07D 209/14							
According to	According to International Patent Classification (IPC) or to both national classification and IPC							
B. FIEL	B. FIELDS SEARCHED							
i	umentation searched (classification system followed l	by classification symbols)						
C07D 209/14	7D 209/14							
Documentatio	n searched other than minimum documentation to the	extent that such documents are included in the	fields searched					
	a base consulted during the intertnational search (nam	e of data base and, where practicable, search ter	ms used)					
stn on the we	≥b .							
C. DOCUM	MENTS CONSIDERED TO BE RELEVANT							
Category*	Citation of document, with indication, where ap	propriate, of the relevant passages	Relevant to claim No.					
X	GB 2345486 A (GLAXO GROUP LTD.), 12 JUL.	2000.	1-3					
	SEE THE WHOLE DOCUMENT.							
X	WO 97/03069 AI (GLAXO GROUP LTD.), 30 JAI	1-3						
	SEE THE WHOLE DOCUMENT							
X	TIDWELL, R. ET AL 'AROMATIC AMIDINES. (1-3					
	BLOCK RESPIRATORY SYNCYTIAL INDUCED PLASMIN, UROKINASE, THROMBIN, AND TR							
	CHEMISTRY (1983), 26(2), PP.294-298.							
x	DE 2731039 A1 (AMERICAN HOME PRODUCTS	S CORP.), 19 JAN. 1978	1-3					
	SEE THE PAGES 1-50							
			_					
			· '					
			•					
	documents are listed in the continuation of Box C.	X See patent family annex.						
	stegories of cited documents: defining the general state of the art which is not considered	"T" later document published after the internation date and not in conflict with the application						
to be of pa	uticular relevence plication or patent but published on or after the international	the principle or theory underlying the inver	ntion					
filing date		considered novel or cannot be considered						
	which may throw doubts on priority claim(s) or which is stablish the publication date of citation or other	step when the document is taken alone "Y" document of particular relevence; the claim	ed invention cannot be					
	ason (as specified) referring to an oral disclosure, use, exhibition or other	considered to involve an inventive step we combined with one or more other such doct	then the document is					
means		being obvious to a person skilled in the art						
	published prior to the international filing date but later iority date claimed	"&" document member of the same patent family	,					
Date of the act	ual completion of the international search	Date of mailing of the international search rep	oort					
28-FEBRUARY 2003-(28:02:2003)		28 FEBRUARY 2003 (28.02.200						
	ling address of the ISA/KR	Authorized officer	60					
9	Korean Intellectual Property Office 20 Dunsan-dong, Seo-gu, Dacjeon 302-701, Republic of Korea	CHO, Hee Won	(影竇)					
63	82-42-472-7140	Telephone No. 82-42-481-5607	写到					

INTERNATIONAL SEARCH REPORT

International application No. PCT/KR02/01843

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
GB 2345486 A	12/07/2000.	GB 9929973A0 ·	09/02/2000
		GB 9915510A0	01/09/1999
WO 97/03069 A	30/01/1997	ZA 9605935A	12/02/1998
		GB 9514265 A0	13/07/1995
		JP 11-508906 T2	03/08/1999
		EP 0843671A1	27/03/1998
		AU 6613996A1	10/02/1997
DE 2731039 A1	19/01/1978	ZA 7703939 A	28/02/1979
		GB 1579678 A	19/01/1980
		AU 507828 B2	28/02/1980
		DK 7703099 A	10/01/1978
		BE 856647 A1	09/01/1978
•		FR 2357563 A1	03/02/1978
		CH 634069 A	14/01/1983

Form PCT/ISA/210 (patent family annex) (July 1998)

This Page is Inserted by IFW Indexing and Scanning Operations and is not part of the Official Record

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images include but are not limited to the items checked:

☐ BLACK BORDERS	
☐ IMAGE CUT OFF AT TOP, BOTTOM OR SIDES	
FADED TEXT OR DRAWING	
☐ BLURRED OR ILLEGIBLE TEXT OR DRAWING	
☐ SKEWED/SLANTED IMAGES	
☐ COLOR OR BLACK AND WHITE PHOTOGRAPHS	
☐ GRAY SCALE DOCUMENTS	
LINES OR MARKS ON ORIGINAL DOCUMENT	
☐ REFERENCE(S) OR EXHIBIT(S) SUBMITTED ARE POOR QUALITY	

IMAGES ARE BEST AVAILABLE COPY.

☐ OTHER:

As rescanning these documents will not correct the image problems checked, please do not report these problems to the IFW Image Problem Mailbox.

THIS PAGE BLANK (USPTO